

Models to Estimate Physical/Chemical Properties of Chemicals

The PHYSICAL/CHEMICAL PROPERTY MODELS included in this section are:

- ❖MPBPVP™
- ❖WSKOWWIN™
- ❖KOWWIN™
- ❖HENRYWIN™
- ❖WVOLWIN™

Following are brief fact sheets providing information on the models OPPT has developed and uses to estimate environmental fate of chemicals. Information provided on each model includes:

- ❖What physical/chemical property does the model estimate?
- ❖What is significant about the physical/chemical property to risk assessment?
- ❖Why is knowing physical/chemical properties important?
- ❖Why would I want to use the model?
- ❖What do I need to run the model?
- ❖What are the inputs and outputs for the model?

MPBPVP™ to Estimate Melting Point, Boiling Point, and Vapor Pressure

What Does the MPBPVP™ Model Do?

MPBPVP™ estimates an organic chemical's melting point, boiling point, and vapor pressure at 25 deg C using a combination of techniques.

Why Is Knowing Melting Point, Boiling Point, and Vapor Pressure Important?

Melting Point (MP), the temperature at which a chemical changes from solid to liquid, gives clues to other chemical properties:

- ❖ MP indicates the state (solid-liquid-gas) of the chemical in the ambient environment.
- ❖ High MP indicates low water solubility.
- ❖ Low MP indicates increased absorption is possible through the skin, GI tract, or lungs.
- ❖ The range of measured MPs indicates its purity: narrow = more pure, wide = less pure.
- ❖ MP < 100°C = increased volatility and higher potential exposures.

Boiling Point (BP), the temperature at which the VP of a chemical in a liquid state equals atmospheric pressure, gives clues to other chemical properties:

- ❖ High BP indicates low VP, for example structurally large substances like polymers.

Vapor Pressure (VP), the pressure at which a liquid and its vapor are in equilibrium at a given temperature, gives clues to other chemical properties:

- ❖ Chemicals with VP $\geq 10^{-4}$ mm Hg (higher VP) exist mostly in the vapor phase, and often have higher potential inhalation exposures than chemicals with low vapor pressure.
- ❖ Chemicals with VP 10^{-5} to 10^{-7} mm Hg exist in both vapor and solids or particulate phases.
- ❖ Chemicals with (lower VP) $\leq 10^{-8}$ mm Hg exist mostly as solids.

Inputs Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs

- ❖ Molecular weight and formula
- ❖ Estimations of melting point, boiling point, and vapor pressure at 25 deg C
- ❖ Chemical structure can be printed or saved as either MDL ISIS SKC file or MDL MOL file

Examples of Melting Point, Boiling Point, and Vapor Pressure Values

Examples of Melting Points at 25 deg C			Examples of Boiling Points at 25 deg C		
CAS RN	Chemical	Degrees C	CAS RN	Chemical	Degrees C
60571	Dieldrin	135	60571	Dieldrin	340
108952	Phenol	-2	108952	Phenol	170
75092	Dichloromethane	-90	75092	Dichloromethane	80
67641	Acetone	-94	67641	Acetone	45
50000	Formaldehyde	-111	50000	Formaldehyde	10

Examples of Vapor Pressures at 25 deg C		
CAS RN	Chemical	Degrees C
60571	Dieldrin	1.77E-5
108952	Phenol	1
75092	Dichloromethane	86
67561	Methanol	396
50000	Formaldehyde	1330

Sample Output from the MPBPVP™ Model

Where Can I Get MPBPVP™?

MPBPVP™ v1.40 has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitedl.htm. If you download and install EPI Suite™ you can run MPBPVP™ as a stand-alone model by putting a shortcut to the MPBPWIN.exe file on your Windows Desktop.

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

INPUTS: CAS Number 108883 (Methyl-benzene or toluene)

RESULTS:

Experimental Database Structure Match:

Name : TOLUENE
 CAS Num : 000108-88-3
 Exp MP (deg C): -94.9
 Exp BP (deg C): 110.6
 Exp VP (mm Hg): 2.84E+01
 Exp VP (deg C): 25
 Exp VP ref : DAUBERT,TE & DANNER,RP (1985)

SMILES : c(cccc1)(c1)C
 CHEM : Methyl-benzene or Toluene
 MOL FOR: C7 H8
 MOL WT : 92.14

----- SUMMARY MPBPWIN v1.40 -----

Boiling Point: 125.72 deg C (Adapted Stein and Brown Method)

Melting Point: -78.09 deg C (Adapted Joback Method)
 Melting Point: -40.26 deg C (Gold and Ogle Method)
 Mean Melt Pt : -59.17 deg C (Joback; Gold,Ogle Methods)
 Selected MP: -59.17 deg C (Mean Value)

Vapor Pressure Estimations (25 deg C):

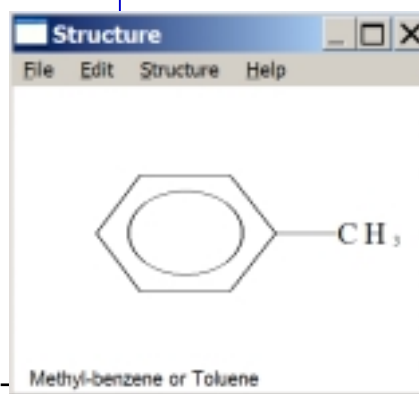
(Using BP: 110.60 deg C (exp database))

(MP not used for liquids)

VP: 25.1 mm Hg (Antoine Method)
 VP: 22.3 mm Hg (Modified Grain Method)
 VP: 29.2 mm Hg (Mackay Method)

Selected VP: 23.7 mm Hg (Mean of Antoine & Grain methods)

TYPE	NUM	BOIL DESCRIPTION	COEFF	VALUE
Group	1	-CH3	21.98	21.98
Group	5	CH (aromatic)	28.53	142.65
Group	1	-C (aromatic)	30.76	30.76
*		Equation Constant		198.18
=====				
RESULT-uncorr		BOILING POINT in deg Kelvin		393.57
RESULT- corr		BOILING POINT in deg Kelvin		398.88
		BOILING POINT in deg C		125.72
TYPE	NUM	MELT DESCRIPTION	COEFF	VALUE
Group	1	-CH3	-5.10	-5.10
Group	5	CH (aromatic)	8.13	40.65
Group	1	-C (aromatic)	37.02	37.02
*		Equation Constant		122.50
=====				
RESULT		MELTING POINT in deg Kelvin		195.07
		MELTING POINT in deg C		-78.09



Melting point is calculated by two different methods, the mean value is determined, and the mean is selected as the melting point.

Vapor pressure is also calculated by different methods, and the mean value is selected as the vapor pressure.

WSKOWWIN™ to Estimate Water Solubility

What Does the WSKOWWIN™ Model Do?

WSKOWWIN™ estimates an octanol-water partition coefficient using the algorithms in the KOWWIN program and estimates a chemical's water solubility from this value. This method uses correction factors to modify the water solubility estimate based on regression against log Kow. This model can tell an assessor if the compound will dissolve in surface water. Water solubility is the degree to which a compound will dissolve in water. It is reported as the amount of the chemical (in milligrams) that will dissolve in 1 liter of water (mg/L).

Why is Knowing Water Solubility Important?

Knowing a chemical's water solubility (WS) is important because this can tell an assessor important information about the chemical's potential risk, for example:

- ❖ Chemicals with low WS:
 - Will have low concentration in aqueous media
 - Have a higher fish BCF
 - Are less likely to be absorbed into mammalian tissues.
- ❖ Chemicals with high WS:
 - Are more likely to be transported along with the water during storm events or to ground water; and
 - Have low log KOW values, and are more likely to be absorbed through GI tract, or lungs. The exception is the case of dispersible molecules like surfactants, and detergents, which can have high predicted log KOWs and can be absorbed through the lung.

Inputs: Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs:

- ❖ Molecular weight and formula
- ❖ Water solubility at 25°C (milligrams per liter)
- ❖ Chemical structure can be printed or saved as either MDL ISIS SKC file or MDL MOL file

Examples of Water Solubility Values

Solubility Classification (mg/L or ppm):

Very soluble	> 10,000
Soluble	> 1,000 - 10,000
Moderately sol.	> 100 - 1,000
Slightly soluble	> 0.1 - 100
Insoluble	< 0.1

Water Solubility Classifications

CAS No.	Chemical	Water Sol. (mg/L)
67561	Methanol	1.00E+06
67641	Acetone	2.20E+05
50000	Formaldehyde	5.74E+04
1912249	Atrazine	2.14E+02
60571	Dieldrin	1.46E-01

Important Notes

WSKOWWIN is not appropriate for dispersible compounds, including surfactants.

Where Can I Get WSKOWWIN™?

WSKOWWIN™ has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitedl.htm. If you download and install EPI Suite™ you can run WSKOWWIN™ as a stand-alone model by putting a shortcut to the WSKOWNT.exe file on your Windows Desktop.

WSKOWWIN™ to Estimate Water Solubility

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

INPUTS: CAS Number 1912249 (atrazine)

RESULTS:

Water Sol: 214.1 mg/L

Experimental Water Solubility Database Match:

Name : ATRAZINE
CAS Num : 001912-24-9
Exp WSol : 34.7 mg/L (26 deg C)
Exp Ref : WARD,TM & WEBER,JB (1968)

SMILES : n(c(nc(n1)NC(C)C)NCC)c1CL

CHEM : Atrazine

MOL FOR: C8 H14 Cl1 N5

MOL WT : 215.69

----- WSKOW v1.40 Results -----

Log Kow (estimated) : 2.82

Log Kow (experimental): 2.61

Cas No: 001912-24-9

Name : Atrazine

Refer : Hansch,C et al. (1995)

Log Kow used by Water solubility estimates: 2.61

Equation Used to Make Water Sol estimate:

$\text{Log } S \text{ (mol/L)} = 0.796 - 0.854 \log \text{Kow} - 0.00728 \text{ MW} + \text{Correction}$
(used when Melting Point NOT available)

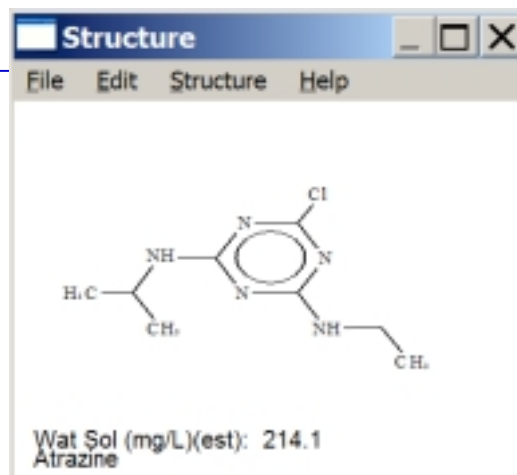
Correction(s):	Value
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No Applicable Correction Factors

Log Water Solubility (in moles/L) : -3.003

Water Solubility at 25 deg C (mg/L): 214.1



KOWWIN™ to Estimate Octanol-Water Partition Coefficient (KOW)

What Does the KOWWIN™ Model Do?

KOWWIN estimates the log octanol-water partition coefficient, log Kow, of chemicals using an atom/fragment contribution method.

Why is Knowing KOW Important?

Knowing KOW will help the risk assessor know where the chemical will go in the stream. KOW indicates whether a chemical predominantly will be found in water (is hydrophilic) or in fatty tissue of animals or other organic materials (is lipophilic) in an aquatic environment. Lipophilic chemicals can bioaccumulate in fatty tissue of fish and bioconcentrate in animals (including humans) that consume the fish. Chemicals with a Log KOW >5 - 6 can bioconcentrate significantly.

Inputs

Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs

- ❖ Log KOW
- ❖ Molecular weight and formula
- ❖ Chemical structure can be printed or saved as either MDL ISIS SKC file or MDL MOL file

Examples of KOW Values

	CAS Number	Chemical	log KOW
lipophilic	25051243	deca-PCB	10.2
	50293	DDT	6.8
	60571	Dieldrin	5.2
	1912249	Atrazine	2.6
	50000	Formaldehyde	0.4
hydrophilic	67641	Acetone	-0.2

log KOW of 0 indicates an equal affinity for lipids and for water

Important Notes

- ❖ KOW is often reported as a **log** due to the extremely wide range of measured KOW values.
- ❖ A log KOW of 0 indicates an equal affinity for lipids and for water.
- ❖ **There is a unique relationship between Log KOW and BCF:** As log KOW increases the solubility in lipids increases. This means an increase in the potential to bioconcentrate in organisms. This relationship begins to change around log KOW of 6. For chemicals with log KOW exceeding 6 the potential to bioconcentrate begins to drop approaching 0 at log KOW of 12.

Where Can I Get KOWWIN™?

KOWWIN™ has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitedl.htm. If you download and install EPI Suite™ you can run KOWWIN™ as a stand-alone model by putting a shortcut to the KOWWINNT.exe file on your Windows Desktop.

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

KOWWIN™ to Estimate Octanol-Water Partition Coefficient (KOW)

Sample Output from the KOWWIN™ Model

INPUTS: CAS Number 60571 (dieldrin)

RESULTS:

Log Kow (version 1.66 estimate): 5.45

Experimental Database Structure Match:

Name : Dieldrin
CAS Num : 000060-57-1
Exp Log P: 5.40
Exp Ref : DeBruijn,J et al. (1989)

Experimental Database Structure Match:

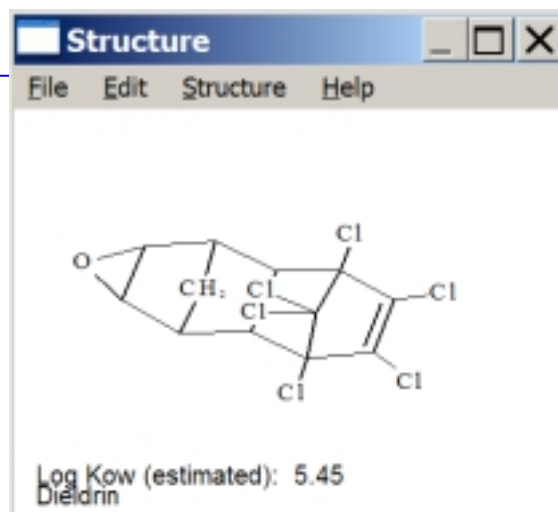
Name : Endrin
CAS Num : 000072-20-8
Exp Log P: 5.20
Exp Ref : DeBruijn,J et al. (1989)

SMILES : CLC4=C(CL)C5(CL)C3C1CC(C2OC12)C3C4(CL)C5(CL)CL

CHEM : Dieldrin

MOL FOR: C12 H8 CL6 O1

MOL WT : 380.91



TYPE	NUM	LOGKOW FRAGMENT DESCRIPTION	COEFF	VALUE
Frag	1	-CH2- [aliphatic carbon]	0.4911	0.4911
Frag	6	-CH [aliphatic carbon]	0.3614	2.1684
Frag	1	C [aliphatic carbon - No H, not tert]	0.9723	0.9723
Frag	2	=CH- or =C< [olefinic carbon]	0.3836	0.7672
Frag	1	-O- [oxygen, aliphatic attach]	-1.2566	-1.2566
Frag	4	-CL [chlorine, aliphatic attach]	0.3102	1.2408
Frag	2	-CL [chlorine, olefinic attach]	0.4923	0.9846
Frag	2	-tert Carbon [3 or more carbon attach]	0.2676	0.5352
Factor	2	Fused aliphatic ring unit correction	-0.3421	-0.6842
Const		Equation Constant		0.2290
Log Kow			=	5.4478

HENRYWIN™ to Estimate Henry's Law Constant

What Does the HENRYWIN™ Model Do?

HENRYWIN™ estimates the Henry's Law Constant (HLC) of an organic compound by two different methods. It also can estimate the HLC of an unknown compound based on the HLC of a known compound. Henry's Law constant (HLC) is the ratio of a chemical's vapor pressure to its water solubility. HLC gives a relative measure of the volatility of a compound from water by measuring the extent to which a compound will partition between water and the air.

Why is Knowing the Henry's Law Constant Important?

Knowing the HLC helps the risk assessor predict the fate of the chemical once it is released to surface water. High HLC indicates chemical is likely to volatilize from solution and partition in air. Low HLC indicates chemical is not likely to volatilize and will remain in surface water.

Inputs

Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs

- ❖ Molecular weight and formula
- ❖ Henry's Law Constant estimated by bond contribution method and by group contribution method (best used for pesticides)
- ❖ Chemical structure can be printed or saved as either MDL ISIS SKC file or MDL MOL file

Examples of Henry's Law Constant Values

CAS Number	Chemical	HLC (atm-m ³ /mole)
75092	Dichloromethane	3.0E-03
50000	Formaldehyde	6.1E-05
67641	Acetone	4.0E-05
67561	Methanol	4.4E-06
60571	Dieldrin	5.4E-07

Volatility Classifications

Very volatile	$\geq 10^{-1}$
Volatile	$10^{-1} - 10^{-3}$
Moderately volatile	$10^{-3} - 10^{-5}$
Slightly volatile	$10^{-5} - 10^{-7}$
Nonvolatile	$< 10^{-7}$

Where Can I Get HENRYWIN™?

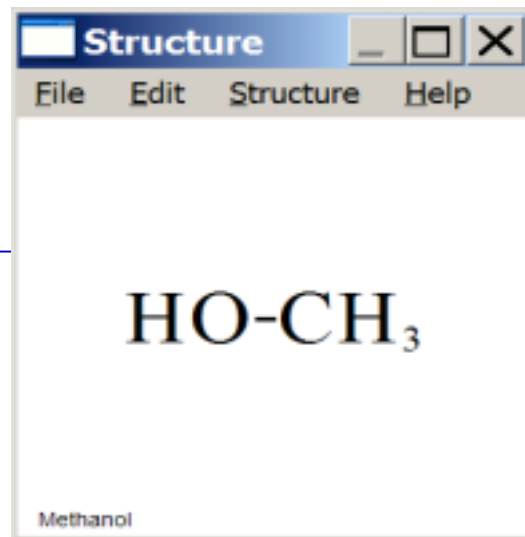
HENRYWIN™ has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitedl.htm. If you download and install EPI Suite™ you can run HENRYWIN™ as a stand-alone model by putting a shortcut to the HENRYNT.exe file on your Windows Desktop.

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

HENRYWIN™ to Estimate Henry's Law Constant

Sample Output From the HENRYWIN™ Model



INPUTS: CAS Number 67561 (methanol)

RESULTS:

Bond Est : 4.27E-006 atm-m3/mole
 Group Est: 3.62E-006 atm-m3/mole

SMILES : OC
 CHEM : Methanol
 MOL FOR: C1 H4 O1
 MOL WT : 32.04

----- HENRYWIN v3.10 Results -----

Experimental Database Structure Match:

Name : METHANOL
 CAS Num : 000067-56-1
 Exp HLC : 4.55E-06 atm-m3/mole
 Temper : 25 deg C
 Exp Ref : GAFFNEY,JS ET AL. (1987)

Two methods are used to estimate HLC. The group contribution method is best used for pesticides.

CLASS	BOND CONTRIBUTION DESCRIPTION	COMMENT	VALUE
HYDROGEN	3 Hydrogen to Carbon (aliphatic) Bonds		-0.3590
HYDROGEN	1 Hydrogen to Oxygen Bonds		3.2318
FRAGMENT	1 C-O		1.0855
FACTOR	* Non-cyclic alkyl or olefinic alcohol		-.2000
RESULT	BOND ESTIMATION METHOD for LWAPC VALUE	TOTAL	3.758

HENRYs LAW CONSTANT at 25 deg C = 4.27E-006 atm-m3/mole
 = 1.74E-004 unitless

	GROUP CONTRIBUTION DESCRIPTION	COMMENT	VALUE
	1 CH3 (X)		-0.62
	1 O-H (C)		4.45
RESULT	GROUP ESTIMATION METHOD for LOG GAMMA VALUE	TOTAL	3.83

HENRYs LAW CONSTANT at 25 deg C = 3.62E-006 atm-m3/mole
 = 1.48E-004 unitless

Models to Estimate Chemical Fate in the Environment

The ENVIRONMENTAL FATE MODELS included in this section are:

- ❖ AOPWIN™
- ❖ HYDROWIN™
- ❖ BIOWIN™
- ❖ PCKOCWIN™
- ❖ BCFWIN™
- ❖ STPWIN™
- ❖ LEV3EPI™

Following are brief fact sheets providing information on the models OPPT has developed and uses to estimate environmental fate of chemicals. Information provided on each model includes:

- ❖ What fate property does the model estimate?
- ❖ What is significant about the fate property to risk assessment?
- ❖ Why is knowing fate properties important?
- ❖ Why would I want to use the model?
- ❖ What do I need to run the model?
- ❖ What are the inputs and outputs for the model?

AOPWIN™ to Estimate Atmospheric Oxidation Potential

What Does the AOPWIN™ Model Do?

AOPWIN™ estimates rate constants and half-lives of atmospheric reactions of organic compounds released to the air with hydroxyl radicals (-OH) and with ozone in the atmosphere. estimates the rate at which certain organic compounds will be destroyed by reactions with compounds in the atmosphere.

Why is Knowing Atmospheric Oxidation Potential Important?

The rate at which an organic compound will be oxidized (destroyed) is an indication of the length of time the compound may reside in the atmosphere.

Inputs

Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs

- ❖ Molecular weight and formula
- ❖ Chemical structure can be printed or saved as either MDL ISIS SKC file or MDL MOL file
- ❖ Hydroxyl radical (-OH) rate constant and half-life
- ❖ Ozone reaction constant and half-life (for olefins and acetylenes only)

Examples of Atmospheric Oxidation Potential Values

CAS Number	Chemical	AOP 1/2 Life (days)
75092	Dichloromethane	79.3
67641	Acetone	52.4
67561	Methanol	17.4
60571	Dieldrin	1.2
1912249	Atrazine	0.4

Atmospheric Oxidation Potential Classifications

Rapid	≤ 2 hrs
Moderate	2 hrs - ≤ 1 day
Slow	> 1 day - ≤ 10 days
Negligible	> 10 days

Important Notes

- ❖ Half-life of >2 days indicates the chemical may be persistent in air.
- ❖ If a chemical has a high AOP rate there still is a potential for inhalation exposure if the travel time from source to receptor is greater than the time for complete oxidation of the compound.

Where Can I Get AOPWIN™?

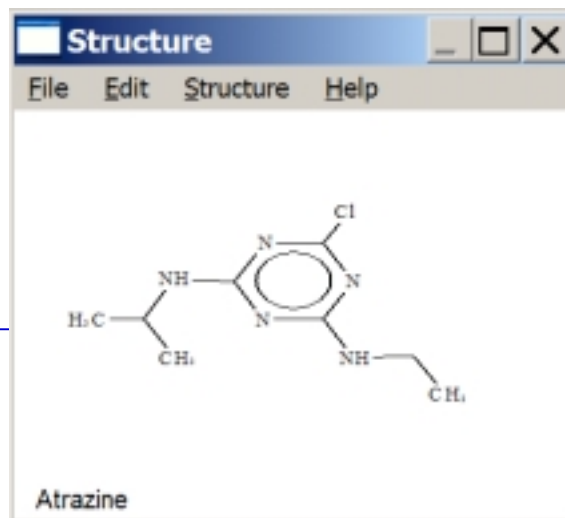
AOPWIN™ has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitd1.htm. If you download and install EPI Suite™ you can run AOPWIN™ as a stand-alone model by putting a shortcut to the AOPWINNT.exe file on your Windows Desktop.

AOPWIN™ to Estimate Atmospheric Oxidation Potential

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

Sample Output from the AOPWIN™ Model



INPUTS: CAS Number 1912249 (atrazine)

RESULTS:

SMILES : n(c(nc(n1)NC(C)C)NCC)c1Cl
 CHEM : Atrazine
 MOL FOR : C8 H14 Cl1 N5
 MOL WT : 215.69

-----SUMMARY : HYDROXYL RADICALS-----

Hydrogen Abstraction	= 24.2300 E-12 cm ³ /molecule-sec
Reaction with N, S, and -OH	= 0.0000 E-12 cm ³ /molecule-sec
Addition to Triple Bonds	= 0.0000 E-12 cm ³ /molecule-sec
Addition to Olefinic Bonds	= 0.0000 E-12 cm ³ /molecule-sec
**Addition to Aromatic Rings	= 0.1176 E-12 cm ³ /molecule-sec
Addition to Fused Rings	= 0.0000 E-12 cm ³ /molecule-sec

OVERALL OH	= 27.3476 E-12 cm ³ /molecule-sec
HALF-LIFE	= 0.391 Days (12-hr day; 1.5E6 OH/cm ³)
HALF-LIFE	= 4.693 Hrs

. . . . * * Designates Estimation(s) Using ASSUMED Value(s)

-----SUMMARY (AOP v1.90): OZONE REACTION-----

* * * * NO OZONE REACTION ESTIMATION * * * *
 (ONLY Olefins and Acetylenes are Estimated)

Experimental Database : NO Structure Matches

Reactions with ozone
are estimated only for
olefins and acetylenes.

HYDROWIN™ to Estimate Hydrolysis

What Does the HYDROWIN™ Model Do?

HYDROWIN™ estimates acid- and base-catalyzed rate constants for chemicals in certain classes (esters, carbamates, epoxides, halomethanes, and certain alkyl halides). A chemical's hydrolytic half-life under typical environmental conditions is also determined. Neutral hydrolysis rates are currently not estimated.

Why is Knowing Hydrolysis Important?

Understanding a chemical's rates of hydrolysis (the breakdown of a chemical by its reaction with water) will help the risk assessor estimate the concentration of the compound in treatment systems and after it is released to surface water, and how long the chemical and/or its hydrolysis products will remain in water bodies after release. The chemical can be catalyzed (broken down) by acids (hydronium) or bases (hydroxide ions).

Inputs

Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs

- ❖ Molecular weight and formula
- ❖ Estimated hydrolysis at 25°C
- ❖ Half-life at pHs 8 and 7
- ❖ Chemical structure can be printed or saved as either MDL ISIS SKC file or MDL MOL file

Examples of Hydrolysis Values

CAS Number	Chemical	Hydrolysis 1/2 Life (yrs)
51796	Carbamic acid, ethyl ester	3326.534
110383	Ethyl decanolate	7.7

Where Can I Get HYDROWIN™?

HYDROWIN™ has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitd1.htm. If you download and install EPI Suite™ you can run HYDROWIN™ as a stand-alone model by putting a shortcut to the HYDRONT.exe file on your Windows Desktop.

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

BIOWIN™ to Estimate Biodegradation

What Does the BIOWIN™ Model Do?

BIOWIN estimates the time required for a compound to biodegrade. Biodegradation is the destruction of a compound by biota, typically microorganisms, in the environment. Primary biodegradation is a change in molecular structure, and ultimate biodegradation is the complete conversion of the molecule to CO₂ and H₂O.

Why is Knowing Biodegradation Important?

Knowing a chemical's biodegradation rate (the time needed for the chemical to partially or completely degrade once it is released to the stream). Knowing the time required for a chemical to be broken down will help the risk assessor estimate the likely degradation in treatment plants and concentration of the chemical at various locations and times after release to surface water. Chemicals with very long biodegradation times may be highly persistent in the environment if they are not subject to destruction by other processes such as photolysis, hydrolysis, etc.

Inputs

Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs

- ❖ Molecular weight and formula
- ❖ Predicted primary and ultimate biodegradation in hours, days, weeks, or months; also predicted, via separate but by linked model, the probability of fast biodegradation using two different methods
- ❖ Chemical structure can be printed or saved as either MDL ISIS SKC file or MDL MOL file

Examples of Biodegradation Values

CAS Number	Chemical	Ultimate Biodeg.
60571	Dieldrin	recalcitrant
1912249	Atrazine	months
75092	Dichloromethane	weeks-months
67641	Acetone	weeks
67561	Methanol	days-weeks

Biodegradation Classifications

Rapid	≥ 60% in	≤ 7 days
Moderate	≥ 30% in	≤ 28 days
Slow	< 30% in	≤ 28 days
Very slow	< 30% in	> 28 days

Important Notes

Two models are used by BIOWIN, a linear and a non-linear regression model. The models are based on regressions against 36 preselected chemical substructures plus molecular weight for experimental biodegradation data for 295 chemicals. The models correctly classified 90% of the chemicals in their training set as rapidly or not rapidly *biodegradable*. Results were slightly better for the nonlinear model.

Where Can I Get BIOWIN™?

BIOWIN™ has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitedi.htm. If you download and install EPI Suite™ you can run BIOWIN™ as a stand-alone model by putting a shortcut to the BIOWINNT.exe file on your Windows Desktop.

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

BIOWIN™ to Estimate Biodegradation

Sample Output from the BIOWIN™ Model

INPUTS: CAS Number = 67561 (methanol)

RESULTS:

SMILES : OC
 CHEM : Methanol
 MOL FOR : C1 H4 O1
 MOL WT : 32.04

----- BIOWIN v4.00 Results -----

Linear Model Prediction : Biodegrades Fast

Non-Linear Model Prediction: Biodegrades Fast

Ultimate Biodegradation Timeframe: Days-Weeks

Primary Biodegradation Timeframe: Days

MITI Linear Model Prediction : Readily Degradable

MITI Non-Linear Model Prediction: Readily Degradable

TYPE	NUM	BIOWIN FRAGMENT DESCRIPTION	COEFF	VALUE
Frag	1	Aliphatic alcohol [-OH]	0.1587	0.1587
MolWt	*	Molecular Weight Parameter		-0.0153
Const	*	Equation Constant		0.7475

RESULT	LINEAR BIODEGRADATION PROBABILITY	0.8910
--------	-----------------------------------	--------

TYPE	NUM	BIOWIN FRAGMENT DESCRIPTION	COEFF	VALUE
Frag	1	Aliphatic alcohol [-OH]	1.1178	1.1178
MolWt	*	Molecular Weight Parameter		-0.4550

RESULT	NON-LINEAR BIODEGRADATION PROBABILITY	0.9752
--------	---------------------------------------	--------

A Probability Greater Than or Equal to 0.5 indicates --> Biodegrades Fast
 A Probability Less Than 0.5 indicates --> Does NOT Biodegrade Fast

TYPE	NUM	BIOWIN FRAGMENT DESCRIPTION	COEFF	VALUE
Frag	1	Aliphatic alcohol [-OH]	0.1600	0.1600
MolWt	*	Molecular Weight Parameter		-0.0708
Const	*	Equation Constant		3.1992

RESULT	SURVEY MODEL - ULTIMATE BIODEGRADATION	3.2883
--------	--	--------

TYPE	NUM	BIOWIN FRAGMENT DESCRIPTION	COEFF	VALUE
Frag	1	Aliphatic alcohol [-OH]	0.1294	0.1294
MolWt	*	Molecular Weight Parameter		-0.0462
Const	*	Equation Constant		3.8477

RESULT	SURVEY MODEL - PRIMARY BIODEGRADATION	3.9310
--------	---------------------------------------	--------

Result Classification: 5.00 -> hours 4.00 -> days 3.00 -> weeks
 (Primary & Ultimate) 2.00 -> months 1.00 -> longer

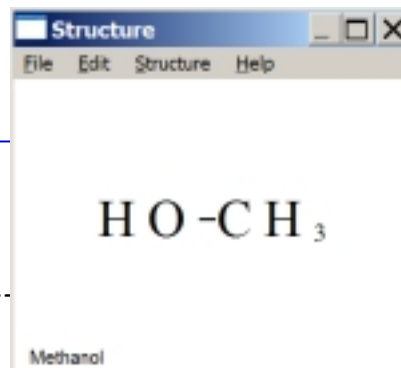
TYPE	NUM	BIOWIN FRAGMENT DESCRIPTION	COEFF	VALUE
Frag	1	Aliphatic alcohol [-OH]	0.1611	0.1611
Frag	1	Methyl [-CH3]	0.0004	0.0004
MolWt	*	Molecular Weight Parameter		-0.0953
Const	*	Equation Constant		0.7121

RESULT	MITI LINEAR BIODEGRADATION PROBABILITY	0.7784
--------	--	--------

TYPE	NUM	BIOWIN FRAGMENT DESCRIPTION	COEFF	VALUE
Frag	1	Aliphatic alcohol [-OH]	1.0041	1.0041
Frag	1	Methyl [-CH3]	0.0194	0.0194
MolWt	*	Molecular Weight Parameter		-0.9250

RESULT	MITI NON-LINEAR BIODEGRADATION PROBABILITY	0.9324
--------	--	--------

A Probability Greater Than or Equal to 0.5 indicates --> Readily Degradable
 A Probability Less Than 0.5 indicates --> NOT Readily Degradable



This chemical is predicted to biodegrade completely in days to weeks.

PCKOCWIN™ to Estimate Organic Carbon Adsorption Coefficient (KOC)

What Does the PCKOCWIN™ Model Do?

PCKOCWIN™ predicts a chemical's soil adsorption coefficient (KOC), which is the ability of a chemical to sorb to the organic portion of soil and sediment. KOC estimations are based on the Sabljic molecular connectivity method with improved correction factors. KOC is the ratio of amount of chemical adsorbed per unit mass of organic carbon (the "OC") in soils, sediments, or sludge to the concentration of the chemical in the solution at equilibrium. KOC indicates whether a chemical is likely to be found in water or the organic carbon portion of soils or sediments.

Why is Knowing the Potential to Sorb to Soil Important?

KOC provides an indication of partitioning to sediments and sludge. Low KOC values suggest the chemical is likely to remain in water, and possibly migrate with ground water, because it is not likely to sorb to soils, sediments, or sludge. High KOC value suggests the chemical will sorb to soils and sediments, or sludge, and is not likely to be found in surface water or migrate with ground water.

Inputs

Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs

- ❖ Estimated KOC
- ❖ Molecular weight and formula
- ❖ Chemical structure can be printed or saved as either MDL ISIS SKC file or MDL MOL file

Examples of KOC Values

CAS Number	Chemical	Log KOC
60571	Dieldrin	4.025
1912249	Atrazine	2.362
75092	Dichloromethane	1.376
106898	Epichlorohydrin	0.652
67641	Acetone	0.297

KOC Classifications

Very strong	≥ 4.5
Strong	3.5 - 4.4
Moderate	2.5 - 3.4
Low	1.5 - 2.4
Negligible	< 1.5

Important Notes

- ❖ Like KOW, KOC is also often reported as a log due to the extremely wide range of measured KOC values.
- ❖ When Log KOC ≥ 4.5 chemical will largely be removed by sorption to sludge in wastewater treatment plants.

Where Can I Get PCKOCWIN™?

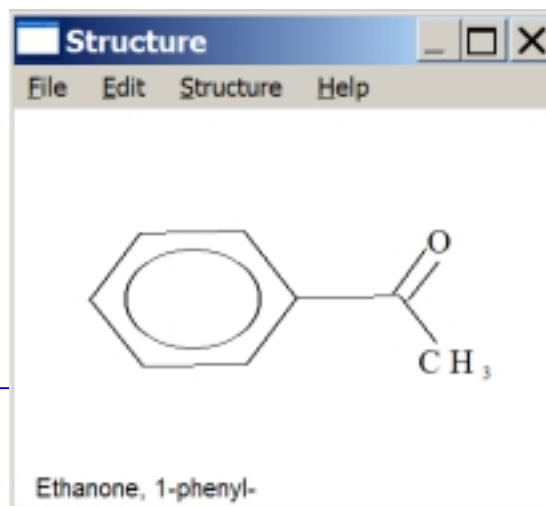
PCKOCWIN™ has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitedl.htm. If you download and install EPI Suite™ you can run PCKOCWIN™ as a stand-alone model by putting a shortcut to the PCKOCNT.exe file on your Windows Desktop.

PCKOCWIN™ to Estimate Organic Carbon Adsorption Coefficient (KOC)

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

Sample Output From the PCKOCWIN™ Model



INPUTS: CAS Number 98-86-2
(Acetophenone)

RESULTS:

Koc (estimated): 46.2

SMILES : O=C(c(ccc1)cl)C
CHEM : Ethanone, 1-phenyl-
MOL FOR: C8 H8 O1
MOL WT : 120.15

----- PCKOCWIN v1.66 Results -----

First Order Molecular Connectivity Index : 4.305
Non-Corrected Log Koc : 2.9123
Fragment Correction(s):
 1 Ketone (-C-CO-C-) : -1.2477
Corrected Log Koc : 1.6646

Estimated Koc: 46.2

BCFWIN™ to Estimate Bioconcentration Factor

What Does the BCFWIN™ Model Do?

BCFWIN™ calculates the BioConcentration Factor and its logarithm from the log Kow. The methodology is analogous to that for WSKOWWIN. Both are based on log Kow and correction factors. A bioconcentration factor (BCF) is the ratio (in L/kg) of a chemical's concentration in the tissue of an aquatic organism to its concentration in the ambient water.

Why is Knowing the Bioconcentration Factor Important?

BCF indicates potential for a chemical to bioaccumulate in lipids (fatty tissue) of aquatic organisms, and to bioconcentrate as it moves up the food web.

Inputs

Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs

- ❖ Estimated Log BCF
- ❖ Molecular weight and formula

Examples of Bioconcentration Factor Values

CAS Number	Chemical	Log BCF
8001352	Toxaphene	4.5
12789036	Chlordane	4.8
60571	Dieldrin	3.7
108703	1,3,5-Trichlorobenzene	2.7

Bioconcentration Factor Classifications

High	≥ 1,000
Moderate	250 - 1,000
Low	< 250

Important Note

There is a unique relationship between Log KOW and BCF: As log KOW increases the solubility in lipids increases. This means an increase in the potential to bioconcentrate in organisms. This relationship begins to change around log KOW of 6. For chemicals with log KOW exceeding 6 the potential to bioconcentrate begins to drop approaching 0 at log KOW of 12.

Where Can I Get BCFWIN™?

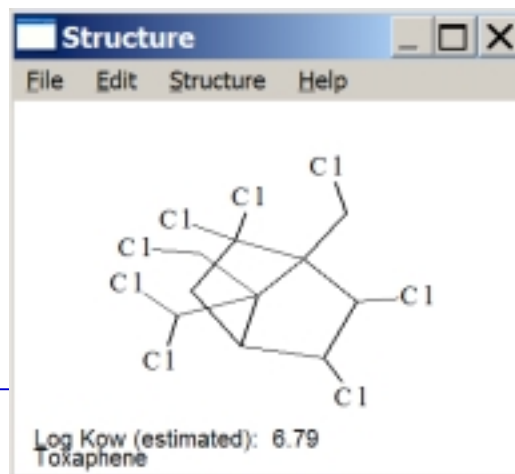
BCFWIN™ has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/EPI_Suitedl.htm. If you download and install EPI Suite™ you can run BCFWIN™ as a stand-alone model by putting a shortcut to the BCFWINNT.exe file on your Windows Desktop.

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

BCFWIN™ to Estimate Bioconcentration Factor

Sample Output From the BCFWIN™ Model



INPUTS: CAS Number 8001352
(toxaphene)

RESULTS:

Log BCF (v2.14 estimate): 3.75

SMILES : CLC(C(CL)C1C2)C(C2(CL)CL)(C1(C(CL)CL)CCL)CCL

CHEM : Toxaphene

MOL FOR: C10 H10 CL8

MOL WT : 413.82

----- Bcfwin v2.14 -----

Log Kow (estimated) : 6.79

Log Kow (experimental): 5.78

Log Kow used by BCF estimates: 5.78

Equation Used to Make BCF estimate:

Log BCF = 0.77 log Kow - 0.70 + Correction

Correction(s):	Value
No Applicable Correction Factors	

Estimated Log BCF = 3.751 (BCF = 5631)

STPWIN™ to Estimate Percent Removal in Wastewater Treatment

What Does the STPWIN™ Model Do?

STPWIN™ predicts the percent of a compound that will be removed from the waste water in wastewater treatment. Values are given for the total removal and three contributing processes (biodegradation, sorption to sludge, and stripping to air) for a standard system and set of operating conditions. STP is "Sewage Treatment Plant" and POTW is "Publicly Owned Treatment Works". Both are names for utilities that treat waste water and usually discharge the treated water to nearby surface water bodies.

Why is Knowing the Percent Removal in Sewage Treatment Plants Important?

Knowing how much of the chemical will be removed from the waste water during wastewater treatment enables the risk assessor to predict how much of the chemical may be discharged by the POTW to surface water and potentially affect aquatic life.

Inputs

Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measure data should be entered as well.

Outputs

- ❖ Estimated percent removal in wastewater treatment
- ❖ Overall chemical mass balance

Examples of Removal Rate Values

CAS Number	Chemical	Percent Removal
60571	Dieldrin	83.11
75092	Dichloromethane	56.91
50000	Formaldehyde	67.3
67641	Acetone	73.06
108952	Phenol	97.47

Where Can I Get STPWIN™?

STPWIN™ has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitedl.htm.

Saving Output

Results can be printed when displayed. After results are displayed click on "Save Results" and you can save results as a ".dat" file that can be opened using MSWord or WordPerfect. Output can also be copied (click on "Copy") through the Windows Clipboard. Structures can be saved as an ISIS ".skc" file or through the Windows Clipboard. Further explanations are given in "Help" on the Results page.

STPWIN™ to Estimate Percent Removal in Wastewater Treatment

Sample Output From the STPWIN™ Model

INPUTS: SMILES : c1ccccc1

RESULTS:

SMILES : c1ccccc1

CHEM : Chemical B

MOL FOR: C6 H6

MOL WT : 78.11

Physical Property Inputs:

Water Solubility (mg/L): 1800

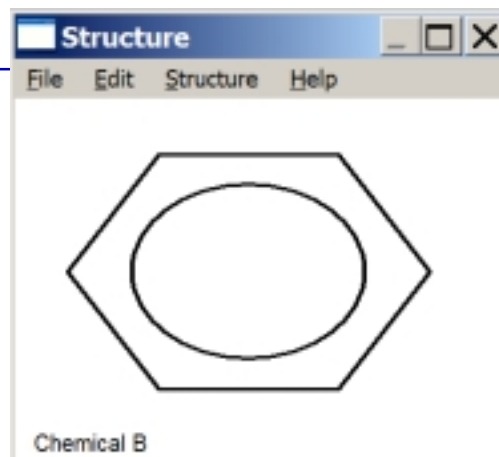
Vapor Pressure (mm Hg) : 95.3

Henry LC (atm-m3/mole) : -----

Log Kow (octanol-water): -----

Boiling Point (deg C) : -----

Melting Point (deg C) : -----



STP Fugacity Model: Predicted Fate in a Wastewater Treatment Facility

=====

PROPERTIES OF: Chemical B

Molecular weight (g/mol) 78.11

Aqueous solubility (mg/l) 1800

Vapour pressure (Pa) 12705.6

(atm) 0.125395

(mm Hg) 95.3

Henry 's law constant (Atm-m3/mol) 0.00555

Air-water partition coefficient 0.226978

Octanol-water partition coefficient (Kow) 134.896

Log Kow 2.13

Biomass to water partition coefficient 27.7793

Temperature [deg C] 25

Biodeg rate constants (h⁻¹), half life in biomass (h) and in 2000 mg/L MLSS (h):

-Primary tank 0.00 526.34 10000.00

-Aeration tank 0.00 526.34 10000.00

-Settling tank 0.00 526.34 10000.00

STP Overall Chemical Mass Balance:

	g/h	mol/h	percent
Influent	1.00E+001	1.3E-001	100.00
Primary sludge	5.63E-002	7.2E-004	0.56
Waste sludge	5.46E-002	7.0E-004	0.55
Primary volatilization	1.25E-001	1.6E-003	1.25
Settling volatilization	1.10E-001	1.4E-003	1.10
Aeration off gas	6.54E+000	8.4E-002	65.43
Primary biodegradation	1.82E-003	2.3E-005	0.02
Settling biodegradation	1.76E-004	2.2E-006	0.00
Aeration biodegradation	2.35E-003	3.0E-005	0.02
Final water effluent	3.11E+000	4.0E-002	31.06
Total removal	6.89E+000	8.8E-002	68.94
Total biodegradation	4.35E-003	5.6E-005	0.04

Predicted
Removal Rate in
Waste Water
Treatment is
69%

LEV3EPI™ Model to Estimate Fugacity

What Does the LEV3EPI™ Model Do?

The LEV3EPI Fugacity model is based on the LEVEL III fugacity model, developed by Don Mackay of Canadian Environmental Modelling Centre, Trent University, Ontario, Canada. LEV3EPI is a non-equilibrium, steady-state multimedia fate model that provides screening level predictions of environmental partitioning (percent in each media) of chemicals between air, soil, sediment, and water under steady state conditions for a default model "environment"; various defaults can be changed by the user. A risk assessor can use this model to estimate where a chemical is most likely to reside once it has been released to the environment.

Why is Knowing a Chemical's Fugacity Important?

If the risk assessor understands a chemical's potential environmental partitioning he/she can use this information to manage the release scenario to minimize detrimental impacts to human health and the environment.

Inputs

Chemical structure (entered as CAS RN and retrieved from the accompanying SMILECAS database; SMILES notation; or drawn and saved as MDL). This program can be operated in a "Batch Mode" so that many structures (as SMILES strings, CAS RNs, or MDL files) can be entered and run at one time. Available measured physical/chemical properties and environmental half-lives should be entered (If not available these properties will be estimated by EPISuite™ components.)

Outputs

- ❖ partition coefficients
- ❖ Z values
- ❖ fugacity of each medium
- ❖ intermedia transport rates and D values
- ❖ reaction and advection D values and loss rates
- ❖ residence times or persistences (overall, reaction, and advection)
- ❖ concentrations and amounts for each medium
- ❖ summary diagram
- ❖ charts of key results, "readme.txt" file with more detailed technical information in a zipped file.

Examples of Fugacity Values

CAS RN	Chemical	Persistence (hrs)
8001352	Toxaphene	4260
12789036	Chlordane	4730
60571	Dieldrin	3040
108703	1,3,5-Trichlorobenzene	791
108952	Phenol	290

Important Notes

Default emission rates are equal amounts (1,000 kg/hr) to air, soil, and water (direct discharges to sediment are unlikely). The model treats a generic environment of 100,000 square km with 10% water; 90% soil surface; water depth 20 m; soil depth 20 cm; sediment depth 5 cm; atmospheric height 1000 m. The LEV3EPI™ model provides a representative environment about the size of the state of Ohio. It is important to note, however, that the percent in each medium will change as a function of the size of the compartments chosen. Moreover, the results of this model are calculated at steady state; a condition that may not occur in the environment (on a global scale).

LEV3EPI™ Model to Estimate Fugacity

Where Can I Get the LEV3EPI™ Model?

The LEV3EPI Fugacity model has been incorporated into the EPI Suite™ which is available at www.epa.gov/opptintr/exposure/docs/episuitedl.htm. If you download and install EPI Suite™ you can run LEV3EPI™ as a stand-alone model by putting a shortcut to the LEV3NT.exe file on your Windows Desktop.

Saving Output

Results can be printed when displayed. After results are displayed click on “Save Results” and you can save results as a “.dat” file that can be opened using MSWord or WordPerfect. Output can also be copied (click on “Copy”) through the Windows Clipboard. Structures can be saved as an ISIS “.skc” file or through the Windows Clipboard. Further explanations are given in “Help” on the Results page.

Sample Output From the LEV3EPI™ Model

Run with default emission rates which assumes equal rates to Air, Water, Soil. Sediment is part of the water column.

Benzene (SMILES c1ccccc1)

	Percent in each media	Half-Life (hrs)
Air	38.0	209
Water	48.0	900
Soil	14.0	900
Sediment	0.2	0

Level III Fugacity Model (Full-Output):

```
=====
Chem Name      : Chemical B
Molecular Wt   : 78.11
Henry's LC     : 0.00555 atm-m3/mole (Henry database)
Vapor Press    : 87.2 mm Hg (MPBPVP™ program)
Log Kow        : 2.13 (Kowwin program)
Soil Koc       : 55.3 (calc by model)
```

	Mass Amount (percent)	Half-Life (hr)	Emissions (kg/hr)
Air	37.6	209	1000
Water	48.4	900	1000
Soil	13.8	900	1000
Sediment	0.202	3.6e+003	0

	Fugacity (atm)	Reaction (kg/hr)	Advection (kg/hr)	Reaction (percent)	Advection (percent)
Air	5.91e-010	627	1.89e+003	20.9	62.9
Water	8.65e-008	188	244	6.25	8.12
Soil	1.64e-007	53.4	0	1.78	0
Sediment	7.75e-008	0.196	0.0203	0.00652	0.000677

```
Persistence Time: 168 hr
Reaction Time:    579 hr
Advection Time:   236 hr
Percent Reacted:  28.9
Percent Advected: 71.1
```

Half-Lives (hr), (based upon Biowin (Ultimate) and Aopwin):

```
Air:      208.7
Water:    900
Soil:     900
Sediment: 3600
Biowin estimate: 2.441 (weeks-months)
```

Advection Times (hr):

```
Air:      100
Water:    1000
Sediment: 5e+004
```

